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(FILE 'HOME' ENTERED AT 16:00:34 ON 19 SEP 2002)

INDEX 'ADISALERTS, ADISINSIGHT, ADISNEWS, AGRICOLA, ANABSTR, AQUASCI,  
BIOBUSINESS, BIOCOMMERCE, BIOSIS, BIOTECHABS, BIOTECHDS, BIOTECHNO,  
CABA,  
CANCERLIT, CAPLUS, CEABA-VTB, CEN, CIN, CONFSCI, CROPB, CROPU, DDFB,  
DDFU, DGENE, DRUGB, DRUGLAUNCH, DRUGMONOG2, ...' ENTERED AT 16:00:41 ON  
19 SEP 2002

SEA (FUSION PROTEIN OR HYBRID PROTEIN OR BI-FUNCTIONAL  
PROTEIN)

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354 FILE ADISALERTS  
124 FILE ADISINSIGHT  
59 FILE ADISNEWS  
1466 FILE AGRICOLA  
91 FILE ANABSTR  
226 FILE AQUASCI  
620 FILE BIOBUSINESS  
513 FILE BIOCOMMERCE  
22793 FILE BIOSIS  
8747 FILE BIOTECHABS  
8747 FILE BIOTECHDS  
19749 FILE BIOTECHNO  
2855 FILE CABA  
18607 FILE CANCERLIT  
31606 FILE CAPLUS  
1124 FILE CEABA-VTB  
29 FILE CEN  
316 FILE CIN  
259 FILE CONFSCI  
51 FILE CROPU  
1 FILE DDFB  
985 FILE DDFU  
55235 FILE DGENE  
1 FILE DRUGB  
215 FILE DRUGNL  
1280 FILE DRUGU  
138 FILE DRUGUPDATES  
358 FILE EMBAL  
20194 FILE EMBASE  
12009 FILE ESBIODASE  
1016 FILE FEDRIP  
43 FILE FROSTI  
345 FILE FSTA  
4817 FILE GENBANK  
8 FILE HEALSAFE  
2764 FILE IFIPAT  
1564 FILE JICST-EPLUS  
7 FILE KOSMET  
12056 FILE LIFESCI  
3 FILE MEDICNF  
48233 FILE MEDLINE  
3 FILE NIOSHTIC  
140 FILE NTIS  
22 FILE OCEAN  
6155 FILE PASCAL

105 FILE PHAR  
49 FILE PHARMAML  
1 FILE PHIC  
213 FILE PHIN  
1303 FILE PROMT  
19183 FILE SCISEARCH  
16814 FILE TOXCENTER  
16804 FILE USPATFULL  
134 FILE USPAT2  
1 FILE VETB  
196 FILE VETU  
4430 FILE WPIDS  
4430 FILE WPINDEX

L1 QUE (FUSION PROTEIN OR HYBRID PROTEIN OR BI-FUNCTIONAL  
PROTEIN)

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FILE 'MEDLINE, BIOSIS, CAPLUS, EMBASE, BIOTECHNO, SCISEARCH' ENTERED AT  
16:04:56 ON 19 SEP 2002

L2 422 S L1 (S) (CELLULOSE BINDING DOMAIN OR CBD)  
L3 43 S L2 (S) ANTIBODY  
L4 12 DUP REM L3 (31 DUPLICATES REMOVED)

=> d 14 ibib ab 1-12

L4 ANSWER 1 OF 12 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 2002:355978 CAPLUS

DOCUMENT NUMBER: 137:105888

TITLE: Use of recombinant cellulose-binding domains of  
Trichoderma reesei cellulase as a selective  
immunocytochemical marker for cellulose in protozoa

AUTHOR(S): Linder, Markus; Winiecka-Krusnell, Jadwiga; Linder,  
Ewert

CORPORATE SOURCE: VTT Biotechnology, Espoo, FIN 02044-VTT, Finland

SOURCE: Applied and Environmental Microbiology (2002), 68(5),  
2503-2508

CODEN: AEMIDF; ISSN: 0099-2240

PUBLISHER: American Society for Microbiology

DOCUMENT TYPE: Journal

LANGUAGE: English

AB Some unicellular organisms are able to encyst as a protective response to  
harmful environment. The cyst wall usually contains chitin as its main  
structural constituent, but in some cases, as in Acanthamoeba, it  
consists

of cellulose instead. Specific cytochem. differentiation between  
cellulose and chitin by microscopy has not been possible, due to the  
similarity of their constituent .beta.-1,4-linked hexose backbones.

Thus,

various fluorescent brightening agents and lectins bind to both cellulose  
and chitin. We have used a recombinant cellulose-binding protein  
consisting of two cellulose-binding domains (CBDs) from Trichoderma

reesei

cellulases linked together in combination with monoclonal anticellulase  
antibodies and anti-mouse Ig fluorescein conjugate to specifically stain  
cellulose in the cysts of Acanthamoeba strains for fluorescence

microscopy

imaging. Staining was obsd. in ruptured cysts and frozen sections of  
cysts but not in intact mature cysts. No staining reaction was obsd.

with

the chitin-contg. cyst walls of Giardia intestinalis, Entamoeba dispar,

or

Pneumocystis carinii. Thus, the recombinant CBD can be used as a marker  
to distinguish between cellulose and chitin. Thirteen of 25

environmental

or clin. isolates of amoebae reacted in the CBD binding assay. All 13  
isolates were identified as Acanthamoeba spp. Five isolates of  
Hartmannella and seven isolates of Naegleria tested neg. in the CBD  
binding assay. Whether cyst wall cellulose really is a unique property

of

Acanthamoeba spp. among free-living amoebae, as suggested our findings,  
remains to be shown in more extensive studies.

REFERENCE COUNT: 39 THERE ARE 39 CITED REFERENCES AVAILABLE FOR  
THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L4 ANSWER 2 OF 12 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 2001:472868 CAPLUS

DOCUMENT NUMBER: 135:78577

TITLE: Method of delivering benefit agent to fabric via  
antibody/fusion protein as binding molecule

INVENTOR(S): Howell, Steven; Little, Julie; Van Der Logt, Cornelis

PATENT ASSIGNEE(S): Paul Erik; Parry, Neil James  
 Unilever N.V., Neth.; Unilever Bc; Hindustan Lever Ltd  
 SOURCE: PCT Int. Appl., 69 pp.  
 CODEN: PIXXD2  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001046356	A2	20010628	WO 2000-EP12529	20001208
WO 2001046356	A3	20020110		

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,  
 CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,  
 HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,  
 LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,  
 SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU,  
 ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,  
 DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,  
 BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

US 2002019324	A1	20020214	US 2000-742693	20001220
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PRIORITY APPLN. INFO.: EP 1999-310431 A 19991222

AB A method of delivering a benefit agent to fabric for exerting a pre-detd.  
 activity useful for stain removal, perfume delivery, and treating collars  
 and cuffs for wear, is provided, wherein the fabric is pre-treated with a  
 multi-specific binding mol. which has a high binding affinity to said  
 fabric through one specificity and is capable of binding to said benefit  
 agent through another specificity, followed by contacting said

pre-treated

fabric with said benefit agent, to enhance said pre-detd. activity to  
 said

fabric. Preferably, the binding mol. is an **antibody** or fragment  
 thereof, or a **fusion protein** comprising a  
**cellulose binding domain** and a domain having a  
 high binding affinity to another ligand which is directed to said benefit  
 agent such as glucose oxidase.

L4 ANSWER 3 OF 12 MEDLINE DUPLICATE 1

ACCESSION NUMBER: 2000275530 MEDLINE

DOCUMENT NUMBER: 20275530 PubMed ID: 10814589

TITLE: Expression, purification and applications of  
 staphylococcal

protein A fused to cellulose-binding domain.

AUTHOR: Shpigel E; Goldlust A; Eshel A; Ber I K; Efroni G; Singer  
 Y; Levy I; Dekel M; Shoseyov O

CORPORATE SOURCE: The Kennedy Leigh Centre for Horticulture Research and The  
 Otto Warburg Center for Agricultural Biotechnology, The  
 Faculty of Agricultural, Food and Environmental Quality  
 Sciences, The Hebrew University of Jerusalem, P.O. Box 12,  
 Rehovot, Israel.

SOURCE: BIOTECHNOLOGY AND APPLIED BIOCHEMISTRY, (2000 Jun) 31 ( Pt  
 3) 197-203.

Journal code: 8609465. ISSN: 0885-4513.

PUB. COUNTRY: ENGLAND: United Kingdom

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200007

ENTRY DATE: Entered STN: 20000810

Last Updated on STN: 20000810

Entered Medline: 20000727

AB Because staphylococcal Protein A (ProtA) binds specifically to IgG, it  
 has

been used for many immunological manipulations, most notably **antibody** purification and diagnostics. Immobilization is required for most of these applications. Here we describe a genetic-engineering approach to immobilizing ProtA on cellulose, by fusing it to **cellulose-binding domain (CBD)** derived from the cellulose-binding Protein A of Clostridium cellulovorans.

The bifunctional **fusion protein** was expressed in Escherichia coli, recovered on a cellulose column and purified by elution at alkaline pH. ProtA-CBD was used to purify IgG from rabbit serum and its ability to bind IgG from different sources was determined. The bifunctional chimaeric protein can bind up to 23.4 mg/ml human IgG at a ratio of 1 mol of ProtA-CBD/2 mol of human IgG, and can purify up to 11.6 mg/ml rabbit IgG from a serum. The ability to bind functionally

active **CBD**-affinity reagents to cellulosic microtitre plates was demonstrated. Our results indicate that a combination of **CBD**-affinity reagents and cellulosic microtitre plates is an attractive diagnostics matrix for the following reasons: (i) cellulose exhibits very low non-specific binding; and (ii) **CBD-fusion proteins** bind directly to cellulose at high density. A unique signal-amplification method was developed based on the ability of ProtA-CBD to link stained cellulose particles to primary **antibody** in a Western blot.

L4 ANSWER 4 OF 12 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 2000:779465 CAPLUS

DOCUMENT NUMBER: 134:38967

TITLE: Phage display of cellulose binding domains for biotechnological application

AUTHOR(S): Benhar, Itai; Tamarkin, Aviva; Marash, Lea; Berdichevsky, Yevgeny; Yaron, Sima; Shoham, Yuval; Lamed, Raphael; Bayer, Edward A.

CORPORATE SOURCE: Department of Molecular Microbiology and Biotechnology, The George S. Wise Faculty of Life Sciences, Tel-Aviv University, Ramat Aviv, Israel

SOURCE: ACS Symposium Series (2000), 769(Glycosyl Hydrolases for Biomass Conversion), 168-189

CODEN: ACSMC8; ISSN: 0097-6156

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

AB A review with 65 refs. In recent years, cellulose-binding domains (CBDs) derived from the cellulolytic systems of cellulose-degrading microorganisms have become a focal point of attention for a wide range of biotechnol. applications. The low cost and availability of cellulose matrixes have rendered CBDs attractive as affinity tags for the purifn. and immobilization of a plethora of proteins. Intensive studies of cellulose degrdn. pathways and the identification of components of the cellulose-degrading machinery have contributed significantly to our understanding of the structure and function of CBDs. The time is now

ripe

to utilize engineered CBDs to improve existing applications and to devise novel ones. Here we describe our recent results of expts. where the Clostridium thermocellum scaffoldin CBD was genetically engineered for such purposes. We describe the development of a novel phage display system where the C. thermocellum **CBD** is displayed as a **fusion protein** with single-chain **antibodies**.

Our system is a filamentous (M13) phage display system that enables the efficient isolation and.

REFERENCE COUNT: 65 THERE ARE 65 CITED REFERENCES AVAILABLE FOR THIS

FORMAT RECORD. ALL CITATIONS AVAILABLE IN THE RE

L4 ANSWER 5 OF 12 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1999:530317 CAPLUS  
 DOCUMENT NUMBER: 131:181955  
 TITLE: Purification of recombinantly prepared proteins by using the cellulose-binding domain of a cellulose-degrading enzyme as an affinity tag  
 INVENTOR(S): Karita, Shuichi; Ohmiya, Kunio; Sakka, Kazuo; Kimura, Tetsuya  
 PATENT ASSIGNEE(S): Toyobo Co., Ltd., Japan  
 SOURCE: Jpn. Kokai Tokkyo Koho, 12 pp.  
 CODEN: JKXXAF  
 DOCUMENT TYPE: Patent  
 LANGUAGE: Japanese  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 11225763	A2	19990824	JP 1998-29410	19980212

AB Purifn. of a protein (enzyme, **antibody**, or hormone) that is expressed as a **fusion protein** with the **cellulose-binding domain (CBD)** of a cellulose-degrading enzyme is described. The fusion protein-contg. cellular ext. is first mixed with an insol., non-crystal cellulose carrier for absorption; the absorbed fusion protein is then eluted with a carbohydrate such as cellobiose, maltose, glucose, or xylose. The target protein is then retrieved from the purifd. fusion protein by digestion with a proteinase such as trypsin. Purifn. of endoglucanase IV of Ruminococcus albus fused with the CBD of xylanase A of Clostridium stercorarium was described.

L4 ANSWER 6 OF 12 MEDLINE DUPLICATE 2  
 ACCESSION NUMBER: 1999160408 MEDLINE  
 DOCUMENT NUMBER: 99160408 PubMed ID: 10049766  
 TITLE: A cellulose-binding domain-fused recombinant human T cell connective tissue-activating peptide-III manifests heparanase activity.  
 AUTHOR: Rechter M; Lider O; Cahalon L; Baharav E; Dekel M; Seigel D; Vlodavsky I; Aingorn H; Cohen I R; Shoseyov O  
 CORPORATE SOURCE: Department of Immunology, The Weizmann Institute of Science, Rehovot, 76100, Israel.  
 SOURCE: BIOCHEMICAL AND BIOPHYSICAL RESEARCH COMMUNICATIONS, (1999 Feb 24) 255 (3) 657-62.  
 Journal code: 0372516. ISSN: 0006-291X.  
 PUB. COUNTRY: United States  
 DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
 LANGUAGE: English  
 FILE SEGMENT: Priority Journals  
 ENTRY MONTH: 199904  
 ENTRY DATE: Entered STN: 19990413  
 Last Updated on STN: 19990413  
 Entered Medline: 19990401

AB The chemokine connective tissue-activating peptide (CTAP)-III, which belongs to the leukocyte-derived growth factor family of mediators, was previously shown to be mitogenic for fibroblasts. However, it has recently been shown that CTAP-III, released from platelets, can act like a heparanase enzyme and degrade heparan sulfate. This suggests that CTAP-III may also function as a proinflammatory mediator. We have successfully cloned CTAP-III from a lambdagt11 cDNA library of PHA-activated human CD4(+) T cells and produced recombinant CTAP-III as a **fusion protein** with a **cellulose-binding domain** moiety. This recombinant CTAP-III exhibited heparanase activity and released degradation products from metabolically labeled, naturally produced extracellular matrix. We have also developed polyclonal

and monoclonal **antibodies**, and these **antibodies**  
against the recombinant CTAP-III detected the CTAP-III molecule in human  
T cells, polymorphonuclear leukocytes, and placental extracts. Thus, our  
study provides tools to examine further immune cell behavior in inflamed  
sites rich with extracellular moieties and proinflammatory mediators.  
Copyright 1999 Academic Press.

L4 ANSWER 7 OF 12 MEDLINE DUPLICATE 3  
ACCESSION NUMBER: 2000014693 MEDLINE  
DOCUMENT NUMBER: 20014693 PubMed ID: 10545273  
TITLE: Matrix-assisted refolding of single-chain Fv- cellulose  
binding domain fusion proteins.  
AUTHOR: Berdichevsky Y; Lamed R; Frenkel D; Gophna U; Bayer E A;  
Yaron S; Shoham Y; Benhar I  
CORPORATE SOURCE: Department of Molecular Microbiology and Biotechnology,  
The George S. Wise Faculty of Life Sciences, Tel-Aviv  
University, Ramat Aviv, 69978, Israel.  
SOURCE: PROTEIN EXPRESSION AND PURIFICATION, (1999 Nov) 17 (2)  
249-59.  
Journal code: 9101496. ISSN: 1046-5928.  
PUB. COUNTRY: United States  
DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
LANGUAGE: English  
FILE SEGMENT: Priority Journals  
ENTRY MONTH: 200001  
ENTRY DATE: Entered STN: 20000124  
Last Updated on STN: 20000124  
Entered Medline: 20000112

AB We describe a method for the isolation of recombinant single-chain  
**antibodies** in a biologically active form. The single-chain  
**antibodies** are fused to a **cellulose binding**  
**domain** as a single-chain protein that accumulates as insoluble  
inclusion bodies upon expression in Escherichia coli. The inclusion  
bodies  
are then solubilized and denatured by an appropriate chaotropic solvent,  
then reversibly immobilized onto a cellulose matrix via specific  
interaction of the matrix with the **cellulose binding**  
**domain** (CBD) moiety. The efficient immobilization that  
minimizes the contact between folding protein molecules, thus preventing  
their aggregation, is facilitated by the robustness of the Clostridium  
thermocellum CBD we use. This CBD is unique in  
retaining its specific cellulose binding capability when solubilized in  
up to 6 M urea, while the proteins fused to it are fully denatured.  
Refolding  
of the **fusion proteins** is induced by reducing with  
time the concentration of the denaturing solvent while in contact with  
the cellulose matrix. The refolded single-chain **antibodies** in their  
native state are then recovered by releasing them from the cellulose  
matrix in high yield of 60% or better, which is threefold or higher than  
the yield obtained by using published refolding protocols to recover the  
same scFvs. The described method should have general applicability for  
the production of many protein-CBD fusions in which the fusion  
partner is insoluble upon expression.  
Copyright 1999 Academic Press.

L4 ANSWER 8 OF 12 MEDLINE DUPLICATE 4  
ACCESSION NUMBER: 2000027007 MEDLINE  
DOCUMENT NUMBER: 20027007 PubMed ID: 10556552  
TITLE: Phage display of a cellulose binding domain from  
Clostridium thermocellum and its application as a tool for  
antibody engineering.



AUTHOR: Berdichevsky Y; Ben-Zeev E; Lamed R; Benhar I  
CORPORATE SOURCE: Department of Molecular Microbiology, The George S. Wise  
Faculty of Life Sciences, Green Building, Room 202,  
Tel-Aviv University, Ramat Aviv 69978, Israel.  
SOURCE: JOURNAL OF IMMUNOLOGICAL METHODS, (1999 Aug 31) 228 (1-2)  
151-62.  
Journal code: 1305440. ISSN: 0022-1759.  
PUB. COUNTRY: Netherlands  
DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
LANGUAGE: English  
FILE SEGMENT: Priority Journals  
ENTRY MONTH: 199912  
ENTRY DATE: Entered STN: 20000113  
Last Updated on STN: 20000113  
Entered Medline: 19991228

AB Phage display of **antibody** fragments has proved to be a powerful tool for the isolation and in vitro evolution of these biologically important molecules. However, the general usefulness of this technology is still limited by some technical difficulties. One of the most debilitating obstacles to the widespread application of the technology is the accumulation of "insert loss" clones in the libraries; phagemid clones from which the DNA encoding part or all of the cloned **antibody** fragment had been deleted. Another difficulty arises when phage technology is applied for cloning hybridoma-derived **antibody** genes, where myeloma derived light chains, irrelevant to the hybridoma's **antibody** specificity may be fortuitously cloned. Here, we report the construction of a novel phage-display system designed to address these problems. In our system a single-chain Fv (scFv) is expressed as an in-frame **fusion protein** with a **cellulose-binding domain (CBD)** derived from the *Clostridium thermocellum* cellulosome. The **CBD** domain serves as an affinity tag allowing rapid phage capture and concentration from crude culture supernatants, and immunological detection of both displaying phage and soluble scFv produced thereof. We demonstrate the utility of our system in solving the technical difficulties described above, and in speeding up the process of scFv isolation from combinatorial **antibody** repertoires.

L4 ANSWER 9 OF 12 MEDLINE DUPLICATE 5  
ACCESSION NUMBER: 1999201269 MEDLINE  
DOCUMENT NUMBER: 99201269 PubMed ID: 10099473  
TITLE: Improved immobilization of fusion proteins via cellulose-binding domains.  
AUTHOR: Linder M; Nevanen T; Soderholm L; Bengs O; Teeri T T  
CORPORATE SOURCE: VTT Biotechnology and Food Research, P.O. Box 1500, FIN-02044, Finland.  
SOURCE: BIOTECHNOLOGY AND BIOENGINEERING, (1998 Dec 5) 60 (5) 642-7.  
Journal code: 7502021. ISSN: 0006-3592.  
PUB. COUNTRY: United States  
DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
LANGUAGE: English  
FILE SEGMENT: Priority Journals  
ENTRY MONTH: 199904  
ENTRY DATE: Entered STN: 19990511  
Last Updated on STN: 19990511  
Entered Medline: 19990426

AB **Cellulose-binding domains (CBDs)** are structurally and functionally independent, noncatalytic modules found in many cellulose or hemicellulose degrading enzymes. Recent biotechnological applications of the **CBDs** include facilitated



protein immobilization on cellulose supports. In some occasions there have been concerns about the stability of the **CBD** driven immobilization. Here we have studied the chromatographic behavior of variants of the *Trichoderma reesei* cellobiohydrolase I **CBD** belonging to family I. Both **CBDs** fused to **antibody** fragments and isolated **CBDs** were studied and compared. Tritium labeling by reductive methylation was used as a sensitive detection method. The **fusion protein** as well as the isolated **CBD** was found to leak from the column at a rate of 0.3-0.5% of the immobilized protein per column volume. However, the leakage could be overcome by using two **CBDs** instead of a single **CBD** for the immobilization. In this way leakage was reduced to less than 0.01% per column volume. The improved immobilization could also be seen as a decreased migration of the protein down the column in extended washes. Copyright 1998 John Wiley & Sons, Inc.

L4 ANSWER 10 OF 12 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS  
INC.DUPLICATE

6  
ACCESSION NUMBER: 1997:109020 BIOSIS  
DOCUMENT NUMBER: PREV199799408223  
TITLE: Comparison of the adsorption properties of a single-chain antibody fragment fused to a fungal or bacterial cellulose-binding domain.  
AUTHOR(S): Reinikainen, Tapani; Takkinen, Kristiina; Teeri, Tuula T. (1)  
CORPORATE SOURCE: (1) VTT Biotechnol. Food Res., P.O. Box 1500, FIN-02044 VTT  
Finland  
SOURCE: Enzyme and Microbial Technology, (1997) Vol. 20, No. 2, pp. 143-149.  
ISSN: 0141-0229.  
DOCUMENT TYPE: Article  
LANGUAGE: English  
AB *Trichoderma reesei* cellobiohydrolase I (CBHI) and *Cellulomonas fimi* cellulase-xylanase (Cex) both have distinct C-terminal **cellulose-binding domains** which belong to different **CBD** sequence families. Two **fusion proteins** comprising a single-chain **antibody** fragment (OxscFv) against 2-phenyloxazolone fused to the two **CBDs** (**CBD**-CBHI or **CBD**-Cex) were constructed. The binding properties of the **fusion proteins** were studied on different cellulosic substrates. It was shown that the **CBD**-Cex binds the **fusion protein** to cellulose more effectively than the **CBD**-CBHI; however, once immobilized, both **fusion proteins** could be eluted from cellulose only with denaturing agents or very low or high pH. Both **fusion proteins** retained equally well their ability to bind the hapten recognized by their **antibody** part.

L4 ANSWER 11 OF 12 MEDLINE DUPLICATE 7  
ACCESSION NUMBER: 97077619 MEDLINE  
DOCUMENT NUMBER: 97077619 PubMed ID: 8920186  
TITLE: Characterization of *Escherichia coli* expressing an Lpp'OmpA(46-159)-PhoA fusion protein localized in the outer membrane.  
AUTHOR: Stathopoulos C; Georgiou G; Earhart C F  
CORPORATE SOURCE: Department of Microbiology, University of Texas, Austin, 78712, USA.  
SOURCE: APPLIED MICROBIOLOGY AND BIOTECHNOLOGY, (1996 Mar) 45 (1-2)

112-9.  
 Journal code: 8406612. ISSN: 0175-75  
 PUB. COUNTRY: GERMANY: Germany, Federal Republic of  
 DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
 LANGUAGE: English  
 FILE SEGMENT: Biotechnology  
 ENTRY MONTH: 199701  
 ENTRY DATE: Entered STN: 19970128  
 Last Updated on STN: 19970128  
 Entered Medline: 19970102

AB The Lpp'OmpA(46-159) **hybrid protein** can serve as an efficient targeting vehicle for localizing a variety of procaryotic and eucaryotic soluble proteins onto the E. coli surface, thus providing a system for several possible biotechnology applications. Here we show that fusion between Lpp'OmpA(46-159) and bacterial alkaline phosphatase (PhoA), a normally periplasmic dimeric enzyme, are also targeted to the outer membrane. However, protease accessibility experiments and immunoelectron microscopy revealed that, unlike other periplasmic proteins, the PhoA domain of these fusions is not exposed on the cell surface in cells having an intact outer membrane. Conditions that affect the formation of disulfide bonds and the folding of the PhoA domain in the periplasm not only did not facilitate targeting to the cell surface but led to lethality when the fusion was expressed from a high-copy-number plasmid. Furthermore, E. coli expressing the Lpp'OmpA(46-159)-PhoA fusion exhibited strain- and temperature-dependent alterations in outer-membrane permeability. Our results are consistent with previous studies with other vehicles indicating that PhoA is not displayed on the surface when fused to cell-surface expression vectors. Presumably, the enzyme rapidly assumes a tightly folded dimeric conformation that cannot be transported across the outer membrane. The large size and quaternary structure of PhoA may define a limitation of the Lpp'OmpA(46-159) fusion system for the display of periplasmic proteins on the cell surface. Alkaline phosphatase is a unique protein among a group of five periplasmic proteins (beta-lactamase, alkaline phosphatase, Cex cellulase Cex **cellulose-binding domain**, and a single-chain Fv **antibody** fragment), which have been tested as passengers for the Lpp'OmpA(46-159) expression system to date, since it was the only protein not displayed on the surface.

L4 ANSWER 12 OF 12 MEDLINE DUPLICATE 8  
 ACCESSION NUMBER: 95133949 MEDLINE  
 DOCUMENT NUMBER: 95133949 PubMed ID: 7832524  
 TITLE: The expression of recombinant proteins on the external surface of Escherichia coli. Biotechnological applications.  
 AUTHOR: Francisco J A; Georgiou G  
 CORPORATE SOURCE: Department of Chemical Engineering, University of Texas at Austin 78712.  
 SOURCE: ANNALS OF THE NEW YORK ACADEMY OF SCIENCES, (1994 Nov 30) 745 372-82. Ref: 37  
 Journal code: 7506858. ISSN: 0077-8923.  
 PUB. COUNTRY: United States  
 DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
 General Review; (REVIEW)  
 (REVIEW, TUTORIAL)  
 LANGUAGE: English  
 FILE SEGMENT: Priority Journals  
 ENTRY MONTH: 199502  
 ENTRY DATE: Entered STN: 19950307  
 Last Updated on STN: 19950307

AB The expression of recombinant proteins on the external surface of Gram-negative bacteria is expected to open the way for a number of significant biotechnological applications, including the development of live bacterial vaccines, the production of whole cell adsorbents, the preparation of whole cell catalysts, and the display and selection of peptide and **antibody** libraries. We have developed a **fusion protein** system for the production of active recombinant proteins on the surface of *Escherichia coli*. Using this system we have expressed beta-lactamase, the *Cellulomonas fimi* exoglucanase Cex as well as its **cellulose binding domain**, and an antidigoxin single chain Fv **antibody** fragment on the cell surface. Recently we have begun to explore some of the potential applications for cell-surface expression.

L6 ANSWER 4 OF 23 CAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 2000:14964 CAPLUS

DOCUMENT NUMBER: 132:83390

TITLE: Topical cosmetic, dermatological, hygienic, or pharmaceutical composition containing antibodies

INVENTOR(S): Breton, Lionel; Pineau, Nathalie; Giacomoni, Paolo

PATENT ASSIGNEE(S): L'Oreal, Fr.

SOURCE: PCT Int. Appl., 40 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000000163	A1	20000106	WO 1999-FR1549	19990628

W: JP, KR, US

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE

FR 2780286	A1	19991231	FR 1998-8341	19980630
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PRIORITY APPLN. INFO.: FR 1998-8341 19980630

AB The invention concerns compns. for topical use, comprising a cosmetic, dermatol., hygienic or pharmaceutical medium, characterized in that it further comprises: at least a first antibody contg. a protein domain recognizing a skin and/or nail and/or lip and/or eyelash epitope, said first antibody being coupled or not with a pigment and/or coloring agent and/or active cosmetic or dermatol. principle; at least a second antibody contg. a protein domain recognizing said first antibody epitope and/or a second antibody contg. a protein domain recognizing a pigment and/or a coloring agent and/or an active cosmetic or dermatol. principle, said second antibodies being coupled or not with a pigment and/or coloring agent and/or an active cosmetic or dermatol. principle. Monoclonal and polyclonal antibodies were prepd. and coupled with a coloring agents. A cosmetic powder contained non-coupled anti(yellow iron oxide) **antibody** 1.6, non-coupled anti(red iron oxide) **antibody** 1.7, non-coupled anti(black iron oxide) **antibody** 1.5, yellow iron oxide 1.6, red iron oxide 1.7, black iron oxide 1.5, Me paraben 0.3, **perfume** 0.2, magnesium carbonate 0.2, talc 73.8, titanium oxide, alumina, glycerin and silica 3, nylon-12 10, dimethicone 2.51, dimethicone and trimethylsiloxaysilicate 0.73, cetyl dimethicone 0.45, and polymethylsilsesquioxane 5 g.